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Choosing Representative Slices of Program Execution for Microarchitecture Simulations: A Preliminary Application to the Data Stream

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Abstract

Microarchitecture simulations are aimed at providing results representative of the behavior of a processor running an application. Due to CPU time constraints, only a few execution slices of a large application can be simulated. The aim of this paper is to propose a technique to choose a few program execution slices representative of the entire execution. We characterize the behavior of each consecutive slice executed. Then we use a statistical classification method to discriminate the execution slices and select the representative ones. In this paper, we detail this approach and apply it to the data stream. Using data cache simulations on the SPEC95 programs, we show that slices representing 1.52 % (average upon all the SPEC95 but one) of the overall program activity are as representative as trace sampling using a 10 % sampling ratio.

Keywords: micro-architecture simulation, trace sampling, classification, data stream characterization, data cache simulation.

1 Introduction

As the complexity of processors is always increasing, microarchitecture simulations using realistic applications consumes always more CPU time. Execution slowdowns to simulate out-of-order execution microprocessors are in the 1,000–10,000 range [3]. However, when exploring new architectural trends, microprocessor architects need their simulations to complete in an affordable amount of time. Therefore, either the target programs are run with reduced input data sets, or the simulations are executed over smaller portions of execution [21, 22, 23]¹. When using reduced input data sets, the simulation results may not accurately

reflect a realistic activity of the processor, so we do not take it into consideration in the remaining of this paper.

To reduce simulation time on a realistic application and input data set, the simulation is often run over an arbitrary fixed number (e.g. a few billions) consecutive executed instructions (a “big slice”). In order to skip the initialization stage, simulation is often disabled during the execution of the first instructions (hundreds of millions, or billions) [4, 27].

Another way to reduce simulation time is the *trace sampling* technique as suggested by [18, 28] for cache simulations. Trace sampling consists in running the simulation over pseudo randomly chosen fixed-size slices of program execution, commonly referred to as *samples* (e.g. samples of 500000 references every 5 million references) or *clusters*².

In both cases, the amount of simulated data is determined by the execution time of the simulation, and/or by previous empirical results (for instance, a 10 % sampling ratio “is known” to give quite good results). Also, for both methods, the representativity of the simulated slices is very questionable.

Commonly used benchmarks have life times of several years (for instance, 3 years for the SPEC92 benchmarks, 5 years for the SPEC95 benchmarks). Microprocessor design projects are also very long efforts spanning over 5 or more years. Then it appears worthwhile spending time, once and for all, to select representative execution slices over which numerous time consuming simulations will be run.

In this paper we propose to use a classification method on several measures gathered over each consecutively executed program slice, in order to select the most representative ones for microarchitecture simulations.

¹Numerous papers in these conference proceedings show the use one of these techniques when performing microarchitecture simulations.

²In the remaining of this paper we only use the term *samples*.

The target program execution is first cut into fixed-size slices (e.g. one million instructions). Then metrics independent from the simulated microarchitecture implementation are applied to the program, and measures are reported for each execution slice.

This provides a characterization of the dynamic program behavior which can be represented by multivariate statistical data (each *individual* is a slice of the execution). Then, we apply a classification method to group execution slices. For each class, we pick out the slice which is the most representative of the class (i.e. the closest to the class center). Finally, the selected slices are weighted by the representativeness of their class among all classes.

Our technique takes part of precise dynamic information gathered on the target programs for selecting representative slices. It also allows us to evaluate *a priori* the representativeness of the selected slices, through the computation of an *indicator*.

In order to validate our approach, we characterized the data stream of the SPEC95 benchmarks to run data cache simulations. However, a characterization of the instruction stream and data dependencies may be added as input to the classification tool to enable complete microprocessor simulations.

In the next section, we discuss related work on program slice selection for microarchitecture simulations. Section 3 develops our approach in detail. Section 4 is an application of the proposed method to the data stream of the SPEC95 benchmarks. Section 5 summarizes this study and presents directions for future development.

2 Related Work

Using reduced traces (or simulating on-the-fly partial program activity) is needed to perform simulations in an affordable amount of time. A great amount of work has been proposed to shorten overall simulation times.

First, trace sampling has been suggested to improve cache [18] or microprocessor [5] simulation speeds. This technique is known in statistics as *cluster sampling* [16]. Several fixed-size trace slices are picked out at regular or (pseudo-)randomly sized intervals and the simulations are run over them. The chosen slices may be more or less representative of the behavior of the target programs, depending on their size and number. Also, simulations results are biased because the state of the simulator at the beginning of a slice is different from the real state produced by full trace simulation. This effect is known as the *cold start effect*.

Similarly, our approach provides a mean to choose execution slices on which to perform simulations. These may also suffer cold start effects but the methods used in trace sampling to reduce it may also apply. In other respect, our approach takes advantage of dynamic program information to derive a representative slice set. In addition, we provide an indicator which gives an idea of the representativity of the selected slice set. This indicator does not need the simulations to be run.

A recent work proposes to use a small (i.e. 50 M. instructions) representative execution slice [25] for micro-architecture simulations. To this end, “*interval*” branch-misprediction data- and instruction-cache miss rates have been measured: i.e. the measures have been computed separately over each million-instruction interval in the target programs. The computed measures clearly exhibit initialization phases of programs. Then, for each program, a 50M. instruction slice is taken after the characterized initialization phase to further run time-consuming simulations. To validate the representativeness of the selected slices, each 50 M-instruction window has been compared to a 250 M-instruction simulation window running a cycle-accurate simulator of an out-of-order execution processor close to an Alpha 21264 [14].

We suspect such a method to be quite risky since the instruction slices are chosen manually and the slice size is arbitrary. Also the validation of the selected slices relies on longer slices which may not be representative of the whole behavior of the target programs (each program considered executed billions of instructions). At the end of subsection 4.4 we compare results from our approach and from a technique close to this, which consists in choosing an instruction slice which size represents 10 % of all the instructions executed. This technique, compared to ours and the trace sampling technique, lead to the more erroneous results.

The representativeness of the reduced trace (or the selected execution slices) when compared to the full original trace (or the complete program execution) is difficult to evaluate *a priori* and necessitates to characterize the behavior of the target program.

Profile-driven sampling [8] was proposed to obtain a representative reduced size trace. This technique uses a fine-grained program profile (frequency of execution of basic blocks) to filter the full trace. In the reduced trace, only a reduced number of occurrences of each executed basic block is kept. This way, statistics such as instruction mix distribution, and basic block size distribution would match those of the full trace.

Profile-driven sampling takes advantage of dynamic program information to derive a reduced trace. However, the trace size reduction is specified as an input parameter of the method: it does not depend on the program characteristics but on the time available for simulation. Instead, the method we propose provides an indicator (see Section 3) which helps in choosing an adequate trace size reduction.

To evaluate the representativeness of a reduced trace against the full trace, Iyengar et al. [13] introduced the *R-metric*. The representativeness of a reduced trace (i.e. the result of the *R-metric* applied to a reduced trace) is an overall score. Based on the contents of the *R-metric*, a heuristic helps in generating reduced traces which are expected to have good *R-metric* scores.

The *R-metric* uses precise information on the basic blocks executed and reduces it to an overall score. With our metrics, we prefer to keep all the measures we gather on the target programs in order not to lose any piece of information for the classification. Also, the *R-metric* depends on microarchitecture implementation details: e.g. branch history table size for branch prediction. In contrast, our metrics for data memory accesses do not, so that they can be used to simulate numerous cache configurations (see 4.1). While the *R-metric* only evaluates a reduced trace *a posteriori*, our approach uses the metrics to *further* reduce the simulated program activity. Ultimately, as indicated by the title of the paper, the *R-metric* can only be used for processor models with infinite cache.

Time varying behavior of the SPEC95 benchmarks are presented in [24]. [24] reports simulation results for these programs each 100 million executed instructions for various modern architectural features: instructions per cycle (IPC), RUU occupancy, cache miss rate, branch prediction miss rate, address prediction miss rate and value prediction miss rate. This study points out that programs have widely different behaviors over time. For this reason, the sections of the program execution which are simulated must be accurately chosen to be representative of the program behavior as a whole. Our method is driven by this key idea, but we feel that program behavior must be characterized by metrics independent from the architectural implementation which is to be simulated. Moreover, our method actually selects program parts to be simulated and evaluates their representativeness.

3 Representative Execution Slice Selection

We propose to use a characterization of the dynamic behavior of the target programs in order to select representative execution slices for microarchitecture simulations. This section presents in detail the process of selecting the slices, and an evaluation of their representativeness.

3.1 Characterization of the Dynamic Program Behavior.

In order to characterize the dynamic behavior, we first divide the program execution into fixed-size slices (e.g. one million instructions). Then metrics are gathered on the program, and reported for each execution slice.

Choosing metrics to characterize this behavior is a difficult issue. For instance, for data cache behavior, one would like to capture the behavior of a large family of caches with distinct parameters such as associativity, size, line size, replacement policy, etc. For the CPU core behavior, one would like to be able to capture the behavior of different numbers of ALUs... We believe that metrics highly related to these behaviors, but independent from the implementation details can be found (e.g. see Section 4 for metrics related to the data stream).

3.2 Actual Slices Selection.

Following commonly used statistics naming, each slice will be called an *observation* or *individual*, and each metric is considered as a set of *variables*. For an observation, each variable is a coordinate in a multi-dimensional space³. This representation of the characterization of the program behavior is a multivariate statistical data [10].

We apply to this multivariate statistical data a classification method to group points which have behaviors close to each others, with respect to the collected metrics. In each group, we retain the most *representative* slice: it is the point closest to the group center.

Simulation results on each chosen slice will be weighted by the representativeness of its class among all classes:

$$\text{Weight}(\text{slice}) = \text{Rep}(\text{class}) = \frac{\text{number of points in the class}}{\text{total number of points}} \quad (1)$$

³For this reason, in the remaining of this paper, we shall also call an observation a (measure) *point*.

3.3 Representativeness of the Selected Slices.

For a given classification, in a class, we define the representativeness of the selected point as the (Euclidean) distance between the selected point and the center of the class. Accordingly, we introduce for the whole “population”, an indicator of the representativeness of all selected points: it is the weighted mean of the representativeness of the selected points in each class (the weights are those of Equation 1).

This indicator, called *wmdc* (weighted mean distances from centers), globally evaluates the representativeness of the selected slices among all execution slices.

4 An Application to the Data Stream

In this section, we present an application of the method described in the previous section to choose representative slices from a characterization of the data stream. We then measure the representativeness of the selected slices by comparing the results of data cache simulations run on all the program slices and run on the selected slices only.

We first detail architecture independent metrics to evaluate the data memory reference stream. Then, we present the classification tool we used for this experiment. Finally, experimental results on the SPEC95 benchmarks are reported and our method is compared to 1) simulations run over an arbitrary big slice, and 2) the systematic statistical trace sampling approach for data cache simulations.

4.1 Metrics Used

Cache memories [26] were introduced to take advantage of the spatial and temporal locality of memory references [12]. The metrics defined below characterize the temporal and spatial locality of data memory accesses for various line sizes in order to simulate a large range of cache configurations.

We first detail the metrics, then in a detailed example we emphasize the intuition behind them.

4.1.1 Measuring Temporal Locality

We evaluate temporal locality by counting the number of executed instructions between two accesses at the same address for each address in the program. This number is the data reuse distance expressed in terms of instructions executed (**RDI**) between two accesses at the same address.

4.1.2 Measuring Spatial Locality

Spatial locality is exploited in caches by the size of the line. In order to catch spatial locality information, we measure the temporal locality we defined above with *several* line sizes⁴. Spatial locality information is then characterized by the *difference* between the temporal locality distributions for several line sizes. Let us illustrate this on the following example.

4.1.3 Example

Let a reference stream be as follows (*a* is an address, numbers are in bytes): *a* (#1), *a* + 4 (#2), *a* + 8 (#3), *a* + 12 (#4), *a* + 16 (#5), *a* + 20 (#6), *a* + 24 (#7), *a* (#8), *a* + 4 (#9), *a* + 8 (#10), *a* + 12 (#11), *a* + 16 (#12), *a* + 20 (#13), *a* + 24 (#14). We assume that these references are done inside loops, so that the number of instruction executed between two references is fixed and equals 10. Note that this reference stream exhibits a good spatial locality.

- If the line size is 4 bytes, then each reference is done at a different line address. All 7 (twice referenced) addresses (i.e. references #8, #9, #10, #11, #12, #13, #14) have a RDI_4 of $10 \times 7 = 70$ instructions. For instance, the line address ' $a + 16 \gg \log_2(4)$ ' is referenced for the second time (#12) 70 instructions after the first time (#5).
- If the line size is 16 bytes, then we assume references to *a*, *a* + 4, *a* + 8, *a* + 12 to match the same line address. Consequently, references to *a* + 16, *a* + 20, *a* + 24 match the same line address. Reference #8 to *a* has an RDI_{16} of $(8 - 5) \times 10 = 30$ instructions; references #9, #10, #11 to *a* + 4, *a* + 8, and *a* + 12 respectively have an RDI_{16} of 10 instructions because they match the same line address as *a*. Reference #12 to *a* + 16 (line address different from the 4 previous references) has an RDI_{16} of $(12 - 7) \times 10 = 50$ instructions...

The distributions of RDI_4 and RDI_{16} are very different, as shown in Figure 1. This difference represents the spatial locality: a 16 byte line takes advantage of the spatial locality, whereas a 4 byte line does not.

If, in contrast, each reference in the reference stream had been really far from the previous ones (poor spatial locality), then the distribution of RDI_{16} would have been the same as the distribution of RDI_4 (each reference would have matched a different line in both cases).

⁴For instance, the data at absolute address '*a*' will be at line address: $a \gg \log_2(\text{line size})$.

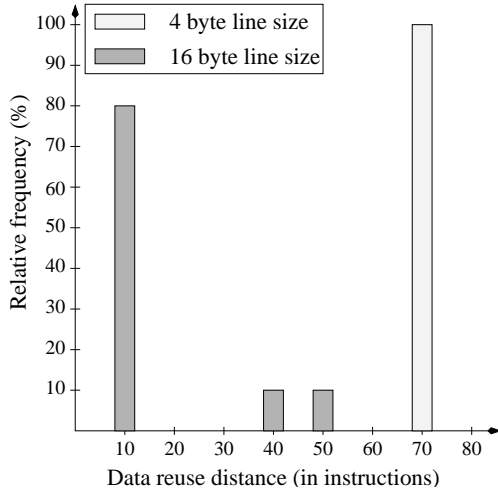


Figure 1: Distributions of data reuse distances for the data stream used in the example (4 byte and 16 byte line sizes).

4.1.4 Data Representation

The multivariate statistical representation of a measured program is composed of individuals (the execution slices) and variables (measures). Given a line size ls , for each slice, and for each integer n , we gather the number of memory accesses which RDI_{ls} is between 2^n and $2^{n+1} - 1$ (this logarithmic scaling limits the number of variables and, accordingly, the amount of data the metric has to produce). For example, the RDI_{16} is represented as follows:

		Variables				
		RDI_{16}				
Individuals	...	$r_{1,1}^{16}$...	$r_{1,j_{16}}^{16}$...	$r_{1,n_{16}}^{16}$
		\vdots		\vdots		\vdots
	...	$r_{i,1}^{16}$...	$r_{i,j_{16}}^{16}$...	$r_{i,n_{16}}^{16}$
		\vdots		\vdots		\vdots
		$r_{N,1}^{16}$...	$r_{N,j_{16}}^{16}$...	$r_{N,n_{16}}^{16}$

The value of the j_{16} th variable for the i th observation ($r_{i,j_{16}}^{16}$) is the reference rate (among all references done during the i th execution slice) which RDI_{16} value is between $2^{j_{16}}$ and $2^{j_{16}+1} - 1$ instructions:

$$r_{i,j_{16}}^{16} = \frac{\text{Card}(\{x \in X_i / RDI_{16}(x) \in [2^{j_{16}}, 2^{j_{16}+1} - 1]\})}{\text{Card}(X_i)}$$

With: $X_i = \{\text{References referenced in the } i\text{th slice}\}$

Note that n_{16} is defined by: $\exists i, r_{i,n_{16}}^{16} \neq 0$, and $\forall i, r_{i,n_{16}+1}^{16} = 0$.

4.1.5 Discussion

The metrics proposed above are very difficult to interpret directly, but they discriminate execution slices against each other, with respect to temporal and spatial locality of data memory accesses. This characteristic is sufficient to apply a statistical classification method.

Moreover, it can be noticed that these metrics do not depend on the slice size: the behavior for a slice, say, twice as big would be similar to the behavior of two smaller consecutive slices. For this reason, the fixed slice size we chose here (see 4.3) does not restrict its generality.

At last, characterizing locality this way have several computational advantages: it is possible to gather several RDI distributions (different line sizes) for all the slices of execution in a single simulation pass. Also, this simulation is far less time consuming than simulating a couple of cache configurations or computing the real LRU distance (i.e. the number of *distinct* references between 2 references at the same (line) address) which necessitates stack processing [2, 15].

4.2 Classification Method Used

Non-hierarchical classification methods [11] are computationally efficient, but they require an *a priori* known number of classes and an estimation of their center location. Instead, hierarchical classification methods [10] “form the final classes by hierarchically grouping subclusters or splitting parent clusters.” For an aggregative (i.e. grouping) method, a hierarchical classification method derives all classification levels from a set of clusters corresponding to the set of points (each point is a cluster), to a unique cluster grouping all the points. The classical representation for a hierarchical classification result is a tree (or *dendogram*), where each level indicates the merging (or splitting) of two subclusters (see Fig. 2).

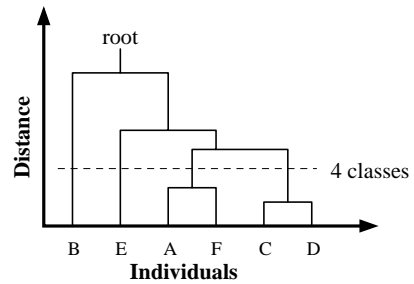


Figure 2: A classification tree.

Once a hierarchical classification method is applied

to a data set, each level in the tree gives a number of classes. Since only one slice (the representative point) will be simulated per class, the amount of simulated data depends on the level chosen in the tree.

We chose to use a hierarchical classification method because it makes it possible to examine all the classification levels before one is chosen. Furthermore, our *wmdc* indicator can be examined for all the levels in the tree.

For this study, we have used a tool called CHAVL which implements the Likelihood Linkage Analysis (LLA) hierarchical classification method [19, 20]. We chose this classification tool because it has proven to be efficient: only a few minutes were necessary to get the classification of a data set containing up to 8000 points with 61 variables. Note that other hierarchical classification tools may be suitable for this study but they may be more time and memory consuming.

4.3 Experimental Setup

We used the **calvin2** + DICE toolset [17] to run the RDI metrics and enable on-the-fly cache simulations. The **calvin2**+DICE toolset consists in:

- an assembly code annotator (**calvin2**) which provides target programs with an efficient fast-forwarding mode,
- and DICE, an embedded emulator which makes it possible to run on-the-fly simulations.

At run time, target programs can switch from direct execution (fast mode) to the emulation mode managed by DICE.

To simulate various level-1 data cache configurations (detailed in Table 1), we connected the Dynero IV on-the-fly cache simulator⁵ to DICE in the target programs.

Varying size (from 4KB to 512KB)
4-way set associative, 32-byte line size, LRU, write back, write allocate.
Varying line size (from 16B to 128B)
4-way set associative, 32 KB, LRU, write back, write allocate.

Table 1: Level-1 data cache configurations simulated.

We tested and validated our approach on the SPEC95 benchmark suite⁶. Each program was

⁵See <http://www.cs.wisc.edu/~markhill/DineroIV/>

⁶Instructions caches were not tested because SPEC95 benchmarks generate too few instruction cache misses [4].

compiled with gcc (version egcs-2.90.29), with the -O3 optimization option, and run on a 143 MHz UltraSPARC-I workstation running Linux (UltraPenguin-1.1.9 distribution, kernel 2.2.10).

For this study, the slice size was not important because we only focussed on the choice of the slices and their representativity among all the consecutive executed slices. For this reason, we chose an arbitrary slice size: one million instructions.

4.4 Experimental Results

For each program execution slice, we collected RDI for line sizes of 16, 64, 256, and 4096 bytes. Experiments showed that this set of line sizes was sufficient to characterize the locality of the data reference stream with respect to the range of data cache configurations we simulated.

Besides, we simulated each SPEC95 benchmark with several level-1 data cache configurations (varying sizes and line sizes) and collected results for every execution slice from the beginning to the end. The purpose of our experiment was not to define a set of representative execution slices that would be used for several years (during the lifetime of a microprocessor design project) since this would have required using many workstations during a month or so. Instead, our experiment only demonstrates the validity of our approach with affordable CPU time. For this reason, on a few applications, we reduced the *train* input data sets to limit the number of executed instructions to a few billions; the other programs were run with their *train* input data set. With these data sets, the number of executed instructions vary from 275 millions to 8 billions, depending on the program. Note that this does not restrict the generality of our results since we evaluated the representativeness of the selected slices among all other slices executed, and since most of the data cache configurations we simulated exhibited non-negligible miss rates.

At last, in order to validate our slice selection approach, for each simulated cache configuration, we compared the weighted mean cache miss rate computed over the selected slices⁷ with the overall mean cache miss rate. To represent this comparison, we computed the relative error:

$$RE(\%) = 100 \times \left| 1 - \frac{\text{weighted mean miss rate (selected slices)}}{\text{overall mean miss rate}} \right|$$

Once, for a given target program, the classification tree has been obtained, there is still a tradeoff to consider to cut the tree. Globally and intuitively, better

⁷The weights are those presented in Section 3.

accuracy in the simulation results is obtained when the tree is cut at a level where many classes appear because many slices are simulated. In other words, the rate of simulated instructions over the entire executed instructions (we call it the *simulation rate*) is high. On the other hand, if the classification tree is cut near its root, a few classes appear, so only a few execution slices are selected and simulated. In this latter case, simulation take less time, but results are expected to be more erroneous.

For the SPEC95 programs, we examined several levels in the classification tree and examined the value of the *wmdc* indicator in conjunction with the *simulation rate*. The slice selection gave accurate results (i.e. relative error smaller than 10 % for absolute cache miss rates higher than 1 %) when the *wmdc* indicator was less than 2 and the simulation rate was more than 0.5 %. For all the programs but *gcc*, obtaining this value of the *wmdc* necessitated to cut the classification tree at level which gave a few classes as summarized in Table 2 (the average relative error is for cache miss rates of more than 1 % among all the cache configurations simulated)⁸.

	<i>wmdc</i>	Sim. rate (%)	Avg. RE (%)
compress95	1.84	1.01	3.56
gcc	1.99	32.41	1.26
go	1.95	0.56	1.56
jpeg	1.97	6.37	0.20
li	0.98	0.56	1.51
m88ksim	0.72	0.40	2.92
vortex	1.02	0.53	2.63
applu	1.59	2.03	1.34
apsi	1.54	0.51	0.26
fp3pp	0.88	0.70	1.16
hydro2d	1.99	3.37	0.17
mgrid	1.54	0.51	2.59
su2cor	1.98	1.51	1.28
swim	1.91	3.66	0.99
tomcatv	1.30	0.51	0.05
turb3d	1.77	0.80	3.82
wave5	1.99	1.33	0.56
Avg.	1.59	3.34	1.52
Avg. (gcc excl.)	1.56	1.52	1.54

Table 2: SPEC95 slice selection: *wmdc* indicator values and average relative errors for trained cache simulations.

Gcc appears to have a characterization difficult to

⁸We did not processed the program *perl* because its cache miss rates appeared to be too low ($\leq 1\%$) for all the cache configurations but one.

classify. Hopefully, our indicator allows us to be aware of this feature. So, we cannot expect to extract a few, very representative slices: we know before running the cache simulations that a simulation rate of less than 32.42 % may lead to erroneous results. However, with a simulation rate of 7.77 % (*wmdc* = 2.63), the maximum relative error for the cache simulation (for cache miss rates less than 1 %) was 5.74 % which is still acceptable despite the relative high value of the *wmdc*. Thus, a high value of the *wmdc* only indicates that we should not be too confident in the representativeness of the simulation results because the average distance of the points from their representant is high.

4.4.1 Comparison with Other Techniques

We compare our method, with a method consisting in choosing one big execution slice after the initialization stage, and the systematic statistical sampling method.

In order to determine the representativeness only of the simulated slices, results for the three methods correspond to cache simulations on the selected slices with trained caches: we used cache simulations on the complete run of the programs, and results were gathered each million simulated instructions.

For the “big slice” method, the resulting cache miss rates are averages upon consecutive 1-million slices representing 10 % of the applications. In order to skip the initialization phase, the slice started in the middle of the execution.

To implement the systematic statistical sampling approach, simulations are run over 1-million instruction samples. The interval between samples is a multiple of 1-million instructions semi-randomly generated to avoid periodic behavior as in [9].

Simulation results with trained caches are displayed in Table 3 (all the numbers are averages for cache miss rates of more than 1 %).

We can notice that both techniques, and particularly the “big slice” one, exhibit high variations between programs. For *m88ksim*, the semi-randomly generated sample set has been exceptionally bad for the 10 % sampling ratio. Note that other sample sets may give better results.

Another results shown by Table 3 is that “big slices” cannot be considered to be representative of overall program behaviors, since the relative errors may or may not be less than 10 % (and globally, they are not).

Comparing Table 3 and Table 2 underlines that our approach with very low simulation rates (except for *gcc*) is globally better than the statistical trace sam-

	Trace sampling		"Big slice"
	5 %	10 %	10 %
compress95	1.59	2.97	9.80
gcc	14.25	2.34	10.79
go	0.37	0.54	5.65
jpeg	12.99	5.06	14.70
li	0.48	0.18	6.11
m88ksim	16.73	130.76	16.73
vortex	12.05	16.17	2.06
applu	3.22	3.20	1.39
apsi	0.76	0.67	1.32
fp3pp	1.18	2.03	0.13
hydro2d	0.48	0.37	2.60
mgrid	2.54	0.35	0.67
su2cor	2.13	0.65	56.98
swim	5.36	4.59	24.49
tomcatv	0.52	0.21	5.26
turb3d	25.47	2.74	66.22
wave5	2.78	1.60	34.47
Avg.	6.05	10.26	15.26
Avg. (m88ksim excl.)	5.39	2.73	15.16

Table 3: Trace sampling and “big slice” technique on the SPEC95: average relative errors for trained cache simulations.

pling technique with a sampling ratio of 10 %. This means that 1) the metrics we chose are well suited to cache simulations, and 2) our approach is really able to extract a few representative execution slices for simulations.

5 Summary and Conclusion

In this paper, we have presented an efficient approach to select representative slices of program execution for microarchitecture simulations. To this end, we first characterize the target program behavior using metrics chosen independent from the further simulated microarchitecture implementation. The gathered program characterization consists of measures for each consecutive fixed size slice of program execution. From this program characterization, we derive a multivariate statistical representation of the program execution (each individual being an execution slice) and then apply a classification method on it. The classification groups sets of “close” execution slices. Then, in each group, the slice nearest to the virtual center is selected (we assume it to be the most representative slice in its group) and weighted by the representativeness of the group (i.e. the number of slices in the group over the total number of slices).

We have applied this method to data memory accesses, and we have compared it with the commonly used technique consisting in choosing one big execution slice and the trace sampling approach. For trained cache simulations on the SPEC95 benchmarks, our method exhibited better results in terms of slice representativity. Also, to obtain results equivalent to the trace sampling method, far less program activity has to be simulated (in most cases less than 1 % is sufficient instead of 10 % for trace sampling).

In this study, we have deliberately focussed on the representativity of the selected slices and therefore reported simulation results for trained caches. However, for real cache simulations, the problem of cold start misses at the beginning of the selected slices is the same as in the trace sampling case, at the beginning of the samples. For this reason, methods applied to the trace sampling approach to reduce this “non-sampling bias” [5] (such as the use of bigger slices, warm-up periods, trace *stitching*... [1, 6, 7, 28]) may also apply to our approach.

Slices selected by our method may be far from the beginning of the program. This necessitates to skip a lot of instructions before the simulation starts as enabled by the direct execution mode of the **calvin2** + DICE toolset [17].

The ultimate use of the method presented in this paper is for complete microprocessor simulations. To this aim, we plan to add several metrics to also characterize the instruction stream (e.g. instruction mix, control transfer instruction characterization), the data dependencies (e.g. distribution of data dependency distances), etc. In the end, a classification will be applied to all the program slices, each of them being characterized by all the metrics.

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